Cell loss from human gastric mucosa measured by the estimation of deoxyribonucleic acid (DNA) in gastric washings

D. N. CROFT, D. J. POLLOCK, AND N. F. COGHILL

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The epithelium of the gastrointestinal tract is constantly being replaced (Bizzozero, 1892, 1893; Friedman, 1945; Bertalanffy and Nagy, 1958) and this is a continuous physiological process (Leblond and Messier, 1958; Creamer, Shorter, and Bamforth, 1961). Gastric surface epithelial and mucus neck cells divide frequently, although the rate of division may be slower in the latter (Stevens and Leblond, 1953). Parietal and chief cells, if they divide at all, do so much more slowly (MacDonald, Trier, and Everett, 1964). In the human stomach surface epithelial cells are formed at the bottom of the gastric pits and migrate up the wall of the pits to reach the surface of the mucosa in two to six days (Lipkin, Sherlock, and Bell, 1963; MacDonald et al., 1964). At the surface they degenerate (Leblond and Stevens, 1948) and are probably shed into the lumen of the stomach in a similar manner to that observed in the small bowel (Stevens and Leblond, 1953; Creamer et al., 1961).

In the resting state an equilibrium should exist between the rate of production and the rate of loss of surface epithelial cells (Stevens and Leblond, 1953; Crosby, 1961). This paper describes a method of measuring the resting rate of accumulation of cells in the gastric lumen which, we suggest, represents the rate of loss of cells from the gastric mucosa. We further suggest that this rate of loss is related to the rate of production of gastric surface epithelial cells.

The cellular content of gastric washings was measured by estimating their deoxyribonucleic acid (DNA) content. As DNA is present in nuclei only and as there is a fairly constant amount in each human somatic cell nucleus (Davidson, Leslie, and White, 1951), the quantity in a gastric washing is proportional to the number of cells present. Previous work in man appeared to show that constant small amounts of DNA were obtained in repeated saline gastric washings (Croft, 1963b). It was thought that such DNA might represent the natural loss of cells from the gastric mucosa. In order to investigate this further the technique for obtaining the gastric washouts was altered and a more sensitive method for estimating DNA (Burton, 1956) was modified to prevent interference from sialic acid present in mucus (Croft and Lubran, 1965).

MATERIALS AND METHODS

GASTRIC PERFUSION TUBE Three holes of 1.0 × 0.3 cm. size were made near the tip of a plastic Ryle’s tube (internal diameter 4.0 mm.). A polythene catheter (internal diameter 2.0 mm.) was attached along the length of this tube so that one end of the catheter opened 9.5 cm. from its uppermost hole (Figure 1.)

REAGENTS Sodium ethylenediamine tetra-acetate (E.D.T.A.) was used to inactivate deoxyribonucleases (Marmur, 1961), and 0.2M E.D.T.A. at pH 8 was added to each washout specimen to give a final concentration of approximately M/100.

Sodium chloride solution (0.9 g./%) used for the washings contained 0.8 mg. polyethylene glycol (M.W. 2

1In receipt of a grant from the Medical Research Council. Present address: St. Thomas' Hospital, London, S.E.1.
4000) per ml. It was intended to use this as an unab-
sorbable marker of infused saline (Hyden, 1965; Borg-
ström, Dahlqvist, Lundh, and Sjövall, 1957) but this
was unnecessary as recovery was virtually complete
in most cases and the application of a correction fac-
tor based on the volume recovered (see below) was sub-
sequently found to be more accurate than one based on
polyethylene glycol recovery. Polyethylene glycol did not
interfere with the chemical estimation of DNA.

Standard DNA solution containing 836 mg atoms
phosphorus per ml was prepared by dissolving 40 mg
of highly polymerized calf thymus DNA (Sigma) in
100 ml 5 mM NaOH and stored at 4°C.

GASTRIC PERFUSION TECHNIQUE Tests were performed in
the morning after the patient had fasted 12 to 15 hours.
Under fluoroscopy the tip of the gastric perfusion tube
was adjusted so that it lay over the left edge of the supine
patient's vertebral column. A dental sucker was placed
in the mouth and the patient asked not to cough, swallow,
or retch during the test. No drugs were given. The pH
of the gastric fluid was measured using narrow range pH
depend. The fasting gastric washout ('first specimen') was
obtained by injecting and withdrawing by syringe 500 ml
of warm (29 to 35°C) saline in 10 50 ml fractions while
rotating the patient to the right and left from the supine
position. Eight of the 50 ml fractions were injected and
withdrawn through the Ryle's tube. In order to wash
out the cardia two fractions were injected down the
polythene catheter and withdrawn through the Ryle's
tube. The time was noted and the polythene catheter at
once attached through a disposable transfusion set to
500 ml of warm saline suspended approximately 80 cm
above the tip of the catheter (Figure 1). The flow of
saline was started by a bellows. The Ryle's tube was
attached through a suction pump (pressure minus 15 to
20 cm. of water) to a litre-sized bottle containing 25 ml
of the 0-2M E.D.T.A. solution. By adjusting the supine
patient's position a continuous return flow was obtained
from the Ryle's tube. If satisfactory flow was not obtained
or if there was bile in the specimen the tube was with-
drawn about an inch. Precautions were taken to prevent
the end of the polythene catheter being withdrawn into
the oesophagus. Saline, 500 ml, was perfused through
the stomach and aspirated over a timed period to yield the
'second specimen'. By the use of a Y-tube connexion
two further bottles of 500 ml of saline were perfused to
give the 'third and fourth specimens'. The specimens
were placed in ice immediately.

CYTOLOGY Specimens for cytological examination were
obtained either during aspiration from a glass sedimenting
chamber attached to the Ryle's tube, or more commonly,
from the particles which had settled to the bottom of the
bottle while being cooled in ice. Particles were floated
on to a slide, fixed in 95% ethyl alcohol, and stained with
haematoxylin and eosin.

RECOVERY OF INFUSED SALINE The average volume re-
covered in the 33 specimens to be reported was 93% of
the volume infused. In three specimens it was less than
80% and in one 112%. Absorption of saline and secretion
of gastric juice could have had little effect on the volumes
recovered because each perfusion took only 10 to 19
minutes (average 13-5) and the perfusing volume was
large (500 ml.).

RATE OF ACCUMULATION OF DNA IN THE STOMACH The
DNA in the whole of each gastric washing was precipi-
tated, extracted, and estimated by the method of
Croft and Lubran (1965). Duplicate estimations differed
by less than ± 3% of their means. The DNA
values were related to the amount of phosphorus in the
standard solution of DNA and were expressed as mg atoms
of DNA-P.

The DNA content of the second, third, and fourth
specimens was corrected for the difference between the
volume of saline infused and the volume of fluid re-
covered by multiplying it by volume infused/volume
recovered. This value was divided by the duration (in
minutes) of the perfusion for the respective specimen, to
yield the rate of accumulation of DNA in the stomach.

GASTRIC DNA RATE The first specimen in all tests con-
tained large amounts of DNA which arose from gastric
and respiratory cells which had accumulated in the
stomach before the start of the test (Croft, 1963b). Much
lower DNA values were found in subsequent specimens,
particularly if the patient did not cough, retch, or swallow.
The lowest rate of accumulation of DNA for each test
was considered to measure DNA loss from the gastric
mucosa and was designated the gastric DNA-rate. Gastric
DNA-rates measured on two occasions separated by five
to 15 weeks on each of four patients were within ±
10% of their means (Table 1).

NON-GASTRIC DNA Contamination of gastric washings
with DNA from material other than cells lost from the
gastric mucosa could have occurred from swallowed
material, duodenal reflux, or DNA-containing bacteria.

### Table 1

<table>
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<th>Patient No.</th>
<th>Clinical Group</th>
<th>Interval between Tests (weeks)</th>
<th>Gastric DNA Rates (mg atoms DNA-P/minute)</th>
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<td>24</td>
<td>Simple atrophic gastritis</td>
<td>15</td>
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Swallowed material To assess the order of contamination from DNA in swallowed saliva and sputum, four patients collected these secretions over timed intervals of 15 to 30 minutes. The DNA content of the material was converted into a secretion rate (Table II). The results indicated that DNA in saliva and sputum, if swallowed, could have increased the lowest gastric DNA-rates by 300% to 1,000%. For this reason a mouth sucker was used and patients with persistent cough and sputum were excluded.

| TABLE II |
| DNA in saliva and sputum from four patients |

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<thead>
<tr>
<th>DNA Secretion Rate (mg. atom DNA-P/min.)</th>
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<tr>
<td><strong>Saliva</strong></td>
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<td>Stimulated with acid drop</td>
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<td><strong>Sputum</strong></td>
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<td>Mucoid (patient with asthma)</td>
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<td>Purulent (patient with pneumonia)</td>
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Cytological examination indicated that the DNA arose mainly from cells lost from the gastric mucosa. Squamous cells were seen in some specimens, and appeared to be more resistant to the action of gastric juice (Tomenius, 1947) than gastric epithelial cells. For this reason squamous cells were probably more conspicuous cytologically than quantitatively important as a source of DNA. As some DNA arose from totally destroyed cells and nuclei and as only a small volume of each 500 ml. specimen was examined, cytology was not an entirely satisfactory method of assessing the amount of contamination with non-gastric DNA.

Duodenal reflux Contamination by duodenal secretions was assessed by noting bile discoloration of the aspirates. This was not present in any specimens from which gastric DNA rates were calculated.

Bacteria Some types of bacteria contain DNA. Bacteria were probably present in the patients with pernicious anaemia (Hurst, 1924; Dick, 1941; Schade, 1960). In our patients with this condition low gastric DNA rates were found which suggested that DNA from bacteria was either quantitatively unimportant, or was removed in the first washing.

The reproducibility of gastric DNA rates even at the lowest values (Table I) suggested that contamination with non-gastric DNA was unimportant. It was unlikely that contamination would have been the same on two separate occasions in each of the four subjects, and Nordgren (1963) prevented contamination of gastric secretions with saliva and duodenal contents by methods similar to those we adopted.

Gastric biopsy Gastric biopsy was usually done within six weeks of the gastric DNA test using a modified Australian gastric biopsy tube (Wood, Doig, Motteram, and Hughes, 1949; Coghill and Williams, 1955). In cases 12, 14, 20, 27, and 28 it was performed between 21 months and 11 years previously. Between one and four specimens (average three) were obtained from each of 27 patients. The biopsy specimens were placed mucosal surface uppermost on duplicating paper and fixed within two minutes in 10% formalin solution. Sections were cut at 6μ and stained with haematoxylin and eosin and by the Feulgen method. Histological examination was made by one observer (D.J.P.) without knowledge of the clinical data or test results. The appearances were classified as follows:—

Normal, including minor miscellaneous mucosal changes (Williams, Edwards, Lewis and Coghill, 1957). Atrophic gastritis, grade I, atrophy of up to 50% of the gastric body glands. Atrophic gastritis, grade II, atrophy of 50 to 90% of the gastric body glands. Atrophic gastritis, grade III, atrophy of 90 to 100% of the gastric body glands.

Particular features of the gastric biopsy specimens are shown in Table III.

Mitosis counts Mitosis counts were performed on the Feulgen-stained gastric biopsy sections, areas in which a gastric gland was cut perpendicularly along its full length being chosen. Epithelial cells were counted from the opening of the gland, down to its lower limit and up the other side to the surface. All mitoses were recorded except early prophase and late telophases. When possible a total of 2,000 cells were counted. The biopsy was generally discarded if, because of bad orientation of the section, less than 1,000 suitable cells could be counted. In some specimens only areas of intestinal metaplasia provided satisfactory sites for counting. There were no differences in the thickness of metaplastic areas in patients with simple atrophic gastritis and pernicious anaemia.

Augmented histamine tests These were performed (Kay, 1953) on patients with pernicious anaemia and on those in whom the pH of a sample of resting gastric juice was above 3·5 using narrow range pH paper. The tip of a Ryle's tube was placed in the stomach by fluoroscopy as described above and 75 mg. of mepyramine maleate was given. The pH of the gastric juice was measured electrometrically.

Gastric antibodies Autoantibody tests were kindly performed by Drs. D. Doniach and I. M. Roitt (Middlesex Hospital). Parietal cell antibodies were detected by immunofluorescence (Taylor, Roitt, Doniach, Couchman, and Shapland, 1962), and the titres determined by complement fixation (Irvine, Davies, Delamore and Williams, 1962; Baur, Roitt, and Doniach, 1965). Antibodies to gastric intrinsic factor were demonstrated by the electrophoretic retention test (Jeffries, Hoskins, and Sleisenger, 1962; Taylor et al., 1962) and by the vitamin B₁₂ charcoal-binding reaction (Ardeman and Chanarin, 1963).

Secretor status Samples of saliva were heated at 100°C. for 10 min., and stored at minus 16 to minus 20°C. Dr. M. J. A. Langman (Guy's Hospital) kindly analysed the specimens for blood group substances (Boyd and Shapleigh, 1954; Clarke, Edwards, Haddock, Howel-Evans, McConnell, and Sheppard, 1956; Doll, Drane, and Newell, 1961).
**TABLE III**

CLINICAL AND LABORATORY DETAILS OF 28 PATIENTS ON WHOM Gastric DNA TESTS WERE PERFORMED

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<th>Case No.</th>
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<th>Clinical Diagnosis</th>
<th>Blood Group</th>
<th>Secretor Status</th>
<th>Parietal Cell</th>
<th>Complementary Fixation</th>
<th>Intrinsic Factor</th>
<th>Gastric Antibodies</th>
<th>Gastric Biopsy</th>
<th>Clinical DNA-rate</th>
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**Group 1: Normal gastric mucosa**

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**Group 2: Treated Addisonian pernicious anaemia**

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**Group 3: Simple atrophic gastritis**

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**SUBJECTS EXAMINED**

The nature of the studies was explained to the patients who agreed to them. The 28 subjects on whom 33 tests were performed fell into three categories (Table III).

**HAEMATOLOGICAL INVESTIGATIONS** Blood group and haemoglobin estimations were performed on all patients. Serum $B_{12}$ concentrations were measured by the method described by Hutner, Bach, and Ross (1956) and radioactive vitamin $B_{12}$ absorption was measured by the faecal method of Booth and Molin (1956), by the method described by Schilling (1953), by the hepatic uptake method (Glass, Boyd, Gellin, and Stephanson, 1954), or by the technique of whole body counting (Belcher, Anderson, and Robinson, 1963).

**PATIENTS WITH NORMAL GASTRIC MUCOSA** Eleven patients (seven males and four females), mean age 44 years (range 29 to 63), had normal gastric mucosa on biopsy. Six of these had dyspepsia with a normal barium meal (non-ulcer dyspepsia), three a duodenal ulcer, one a peptic ulcer and one a smaller lesser curve gastric ulcer. In all, the pH of resting or post-histamine gastric juice was below 3.5. All had a haemoglobin above 12.0 g%. Most of the patients were free of symptoms at the time of the gastric DNA test.

**PATIENTS WITH PERNICIOUS ANAEMIA** Nine patients
(five males and four females), mean age 58 years (range 40 to 72), had Addisonian pernicious anaemia. The pre-treatment serum $B_{12}$ level was below 60 µg/ml, in the eight patients in whom it was estimated. In all nine radioactive vitamin $B_{12}$ absorption was within the range found in Addisonian pernicious anaemia when the dose was given alone or with carbachol and rose to normal levels when the dose was given with intrinsic factor. The histological features of the gastric mucosa are shown in Table III. The gastric biopsy tube could not be passed in case 19 who was achlorhydric. In six patients the pH of the gastric juice rose after maximal histamine stimulation, and in all cases was between 6-8 and 8-0. Case 18 had a tubeless augmented histamine test using Diagnex blue (Squibb); no dye was excreted in the urine, indicating achlorhydria. Five of the patients (cases 12, 14, 16, 18, and 20) had a barium meal and in each the appearances of gastric atrophy were seen (Laws and Pitman, 1960) Gastric DNA tests were performed on one patient (case 19) before treatment and on all when they had been treated with vitamin $B_{12}$ for six weeks to 11 years, and their haemoglobin was over 11.0 g. %.

**PATIENTS WITH SIMPLE ATROPHIC GASTRITIS** Eight patients (five males and three females), mean age 53 years (range 36 to 75), had simple atrophic gastritis (grades I to III) diagnosed by gastric biopsy (Table III). These patients did not have pernicious anaemia and had no family history of it. Five of them had suffered from hypochromic anaemia and three from non-ulcer dyspepsia. Four of these patients secreted gastric HCl (Table III). Of the other four, in two the pH rose, in one it remained the same (6-9), and in one it fell from 5-8 to 5-3 after histamine. The serum $B_{12}$ concentration was 170 µg/ml or above in seven of the patients. Measurements of radioactive vitamin $B_{12}$ absorption in cases 21, 25, 27, and 28 indicated that they were not suffering from pernicious anaemia; case 26 was not tested but secreted gastric HCl. One patient (case 22) with hypochromic anaemia (corrected by oral iron without the development of macrocytosis), had a gastric juice pH of 7-0 after maximal histamine stimulation, grade III atrophic gastritis, and a serum $B_{12}$ level of 95 µg/ml. This patient (tested by the method of whole body counting) absorbed 0-14 µg of an oral dose of 1 µg radio- $B_{12}$ given alone, 0-27 µg. when the dose was given with Carbachol, and 0-47 µg. when the dose was given with intrinsic factor; he resembled patients with simple atrophic gastritis described by Wood, Ralston, Ungar, and Cowling (1964) and by Whiteside, Mollin, Coghill, Williams, and Anderson (1964), and has not developed pernicious anaemia during 18 months of observation. All but one (case 21) had a barium meal. In four it was normal; in three (cases 22, 23, and 25) it showed the appearances of gastric atrophy (Laws and Pitman, 1960). At the time of the gastric DNA test the haemoglobin in these patients was more than 10-0 g. %.

**RESULTS**

**GASTRIC DNA-RATES IN THE THREE CLINICAL GROUPS** The results of 28 gastric DNA-rate determinations, four of which were the mean of duplicate tests, are shown in Fig. 2 and Table III. The rates appeared to be unrelated to age, sex, or duration of disease. Two broad patterns emerged:—

1. Most of the patients with a normal gastric mucosa, or with treated pernicious anaemia, had gastric DNA-rates of 18 mµg. atoms DNA-P per minute or less. Only one patient in each of these groups had a rate above this value.

2. The DNA-rates of patients with simple atrophic gastritis had a wider scatter and in most were higher, only one patient having a rate below 18 µmg. atoms DNA-P per minute. Case 25 had been studied previously by a gastric washout technique involving manual suction and on that occasion had also been found to have a high resting rate of accumulation of DNA (Croft, 1963a).

**EFFECT OF VITAMIN $B_{12}$ ON GASTRIC DNA** While developing the present technique tests which differed from the final method were performed on patients with pernicious anaemia before and up to seven weeks after starting treatment with vitamin $B_{12}$. It appeared that more DNA was obtained in gastric washings after two or three weeks' treatment with vitamin $B_{12}$ than was obtained before, or up to a week after, starting treatment. Gastric DNA tests
1 In those with a normal mucosa denuded nuclei were usually seen. These were considered to come from the gastric surface epithelial cells because in some areas they were present with degenerate but recognizable gastric columnar cells (Figure 4). Intact squamous cells were not uncommon but although cytologically well preserved, they were relatively few in number compared with the denuded nuclei. Polymorphonuclear leucocytes were sometimes present but were not conspicuous.

2 In the patients with treated pernicious anaemia intact gastric surface epithelial cells were seen more commonly. These cells were usually columnar in shape and denuded nuclei were also present. In a few specimens rounded cells of variable size with clumped nuclear chromatin were seen. These had the appearance of abnormal gastric surface epithelial cells (Rubin, 1955; Gibbs, 1962 and 1964;
Numerous polymorphs were present in association with the abnormal gastric surface epithelial cells.

(b) The appearances of the cellular particles in the three patients (cases 21-23) with a relatively low gastric DNA-rate (below 40 mμg. atoms DNA-P per minute) resembled those of the patients with pernicious anaemia. The specimens contained mainly intact columnar cells and denuded gastric epithelial nuclei. There were a few inflammatory cells and some abnormal gastric surface epithelial cells with nuclear clumping.

These results indicated that both in patients with a normal gastric mucosa and those with treated pernicious anaemia the DNA arose mainly from exfoliated gastric surface epithelial cells. These cells were intact in patients with pernicious anaemia, and denuded of cytoplasm in patients with a normal gastric mucosa (and presumably normal gastric peptic activity). In patients with simple atrophic gastritis the DNA was derived from a combination of gastric surface epithelial cells and inflammatory cells extruded from the gastric mucosa (see below).

**GASTRIC HISTOLOGY, DNA-RATES, AND CYTOLOGY**

High DNA-rates (above 40 mμg. atoms DNA-P per min.) were found in the simple atrophic gastritis patients with grades I, II, and III atrophic gastritis. Inflammatory cells in the superficial and deep mucosa were seen more commonly in the biopsy specimens from patients with simple atrophic gastritis or pernicious anaemia than in those of the control group. Inflammatory cells, usually neutrophils, penetrating the surface epithelium were observed in the biopsy specimens of two patients with pernicious anaemia and in specimens from all but one of the patients with simple atrophic gastritis (Figures 6 and 7). The single exception in the last group was case 21, the one patient with simple atrophic gastritis, who had a very low gastric DNA-rate (7 mμg. DNA-P per min.). Thus inflammatory cells were found in most cases of simple atrophic gastritis both in the gastric biopsy specimens and in the cytological preparations, and they were associated with DNA-rates higher than those in most of the patients with a normal mucosa or pernicious anaemia.

The surface epithelium was composed of regular columnar cells in all the patients of the first two groups (Table III), and in four of those with simple atrophic gastritis. In the other four patients with this condition (cases 24-26 and 28) the surface epithelium was composed wholly, or in part (one patient), of irregular cuboidal cells and these patients were among the five with both the highest DNA-rates in this group and numerous abnormal

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**FIG. 5.** Round cells of variable size containing nuclei with clumped chromatin from a perfusion test on a patient with simple atrophic gastritis and high gastric DNA-rate (case 25). These cells were considered to be abnormal gastric surface epithelial cells (haematoxylin and eosin, × 1,600).

Nieburgs and Glass, 1963) and were similar to those seen in patients with simple atrophic gastritis (Figure 5). Polymorphonuclear leucocytes were seen but were not more obvious than in the group with normal gastric mucosa.

3 In the patients with simple atrophic gastritis two patterns were observed.

(a) In the five patients (cases 24-28) with a high gastric DNA-rate (above 40 mμg. atoms DNA-P per minute) both the gastric washings and the slide preparations were macroscopically more cellular. Intact circular cells with nuclei of variable size and containing chromatin clumps were present in large numbers (Fig. 5). These cells were similar to the abnormal gastric surface epithelial cells seen in some specimens from the patients with pernicious anaemia but were more numerous. In some specimens columnar cells were seen but these were relatively few.
gastric surface epithelial cells in the cytological preparations.

MITOSIS COUNTS OF THE THREE CLINICAL GROUPS
These are shown in Table III and Figure 8. In the group with a normal mucosa there was a mean of 0.63 mitoses per 100 surface gastric epithelial cells (S.D. 0.19); in the group with pernicious anaemia there was a mean of 1.4 mitoses (S.D. 0.42) which was significantly higher (P <0.001); in the group with simple atrophic gastritis there was a mean of 1.5 mitoses (S.D. 0.63) which was also significantly higher than in the group with normal mucosa (P <0.001). Three of the four highest mitosis counts were in areas of intestinal metaplasia (two patients with pernicious anaemia and one with simple atrophic gastritis).

SERUM GASTRIC ANTIBODIES None of the patients with normal gastric mucosa had either parietal cell or intrinsic factor antibodies (Table III). Of the nine pernicious anaemia patients all reacted with parietal cells, and six had intrinsic factor antibodies.

FIG. 6. Gastric biopsy specimen from case 23 showing polymorphonuclear neutrophil cells passing into the surface epithelium of the gastric mucosa (Gibbs, 1964) (haematoxylin and eosin, × 1,290).

FIG. 7. The presence or absence of inflammatory cells penetrating the surface epithelium in gastric biopsy specimens from eleven patients with a normal mucosa, eight with treated pernicious anaemia, and eight with simple atrophic gastritis.

FIG. 8. The distribution of mitosis counts in 22 patients from whom satisfactory gastric biopsy specimens were obtained. ○, mitosis count from area of gastric type epithelium. ●, mitosis count from area of intestinal metaplasia. ††, mean of mitosis counts obtained from an area of gastric type epithelium and an area of intestinal metaplasia.
in addition to the normal incidence (Roitt, Doniach, and Shapland, 1965) Of the eight patients with simple atrophic gastritis three had parietal cell antibodies and none reacted against intrinsic factor. There was no correlation between DNA-rates and the presence or absence of gastric antibodies.

SECRETOR STATUS AND BLOOD GROUP All but four patients were secretors of H substance. The gastric DNA-rates of the four non-secretors ranged from low to high values (Table III). In this small series there was no correlation between gastric DNA-rate and blood group or secretor status.

DISCUSSION

Our results have established that the gastric DNA-rate gave a measure of cell loss from the gastric mucosa. There was a higher rate of cell loss in most patients with simple atrophic gastritis than in most of those with a normal mucosa, or with pernicious anaemia. There was a higher rate of production of surface epithelial cells, as judged by mitosis counts, in patients with the atrophic mucosa of pernicious anaemia or simple atrophic gastritis. In the one patient with pernicious anaemia tested the DNA-rate increased significantly after treatment with vitamin B₁₂.

New gastric surface epithelial cells are being constantly formed in the bases of the gastric pits. This continuous production of cells is balanced by an equal rate of destruction (Stevens and Leblond, 1953; Crosby, 1961). As the result of an equilibrium between production and loss the histological appearance of the tissue remains unaltered. Epithelial cells in the gut are destroyed by exfoliation into its lumen and it has been calculated that in this process half a pound of cells is lost from the human gastrointestinal surface per day (Leblond and Walker, 1956). It seems probable that DNA in the nuclei of degenerating epithelial cells is also extruded into the gut lumen, although recent work suggests that some of the DNA from degenerating cells may be re-utilized in the epithelium by newly dividing cells (Cutright, 1965). From data on the DNA content of human cells (Davidson et al., 1951) it can be calculated that the rate of loss of cells in our patients with a normal gastric mucosa ranged from 0·26 to 1·75 million cells per minute with a mean of 0·55 million per minute. When related to unit surface area this order of cell loss corresponds reasonably well with that calculated on theoretical grounds for the human small intestine which is a similarly dynamic tissue (Crosby, 1961).

We have found that measurements of the gastric DNA-rate were the same on different occasions in the same patient and our cytological evidence suggests that the gastric DNA-rate represents the rate of loss of cells from the gastric mucosa. In patients with a normal mucosa and those with pernicious anaemia the exfoliated cells seemed to be mainly intact or denuded gastric surface epithelial cells. In these patients, if it is assumed that the rate of surface epithelial cell loss is in equilibrium with the rate of production of these cells, measurement of gastric DNA-rate should be an index of the turnover of gastric surface epithelium. In the case of most of the patients with simple atrophic gastritis, however, gastric DNA-rates measured loss of both gastric epithelial cells and inflammatory cells extruded from the gastric mucosa. In this group, therefore, the gastric DNA-rate may not have been as satisfactory a measure of surface epithelial cell loss as in the other two groups, although for reasons stated below, it may still have been an index of surface epithelial cell turnover.

Vitamin B₁₂ deficiency affects the synthesis of nucleic acids (Rose and Schweigert, 1952; Smith, 1965) and it has been suggested that this causes a decrease in nucleic acid production and fall in the rate of cell division (Arnstein, 1955). Epithelial tissues in which there is a high rate of cell division might be expected to be especially affected by B₁₂ deficiency, and the glossitis of untreated pernicious anaemia which is corrected by B₁₂ replacement is probably an example of this (Stone and Spies, 1948). Alterations have been reported in the appearance of the nuclei of exfoliated gastric surface epithelial cells from patients with untreated pernicious anaemia (Graham and Rheault, 1954; Raskin, Kirsner, and Palmer, 1958; Schade, 1960) which suggests that vitamin B₁₂ deficiency affects these cells also. The three-fold increase in the rate of accumulation of gastric DNA that was observed in the patient with pernicious anaemia after treatment with vitamin B₁₂ (Fig. 3) could be a reflection of an increased rate of turnover of gastric surface epithelial cells. The histological appearance of the atrophic gastric mucosa of pernicious anaemia is considered not to alter materially after treatment with vitamin B₁₂ so that if there is an increased rate of production of gastric epithelial cells after treatment then there must be a proportional increase in the rate of cell loss. This concept would be in accord with the finding in rat and human small intestine that alteration of surface epithelial cell production (or turnover) does not necessarily lead to changes in the length or shape of the intestinal villi (Trier, 1962; Conrad, Weinstaub, Merrill, and Crosby, 1965).

In normal adults there is only a four-fold range in gastric size and macroscopic surface area (Cox, 1952). Therefore variations in these were unlikely to have accounted for the twenty-fold difference...
between the lowest and highest gastric DNA-rates in our patients. Card and Marks (1960) have estimated total mucosal volume and glandular mucosal volume in partial gastrectomy specimens and from their data an estimate of mucosal epithelial volume can be made by calculation. This figure is probably an index of the number of surface epithelial cells in a stomach and can be shown from their data to have a linear relationship to the maximum acid output over a thirteen-fold range. This implies that there are fewer surface epithelial cells in achlorhydic than in acid-producing stomachs with a normal mucosa, and it is therefore reasonable to assume that there are fewer of these cells in atrophic than in normal stomachs. As the DNA-rates of the group with pernicious anaemia were much the same as those of the patients with a normal mucosa it could be argued that, per unit of gastric surface epithelium, the loss of DNA was higher from these atrophic mucosae.

Although some of the patients with simple atrophic gastritis were not achlorhydic they probably also possessed fewer than normal gastric epithelial cells. It was in this group that in addition to inflammatory cells numerous abnormal surface epithelial cells were noted both in the gastric washings and in the biopsy specimens. Two of the patients with simple atrophic gastritis, who had inflammatory cells penetrating the surface epithelium, and the only two patients with pernicious anaemia showing this phenomenon, had low or relatively low gastric DNA-rates. This, coupled with the cytological appearances, suggested that the high gastric DNA-rates in simple atrophic gastritis were due not only to loss of inflammatory cells but also to high rates of loss of surface epithelial cells. Therefore we conclude that in the majority of patients with simple atrophic gastritis or the atrophic gastritis of pernicious anaemia the gastric DNA-rates indicated that per unit of gastric epithelium there was a higher turnover of surface epithelial cells than in the patients with a normal gastric mucosa. This conclusion is supported by the finding of higher mitosis rates, which usually (but not always) mean higher rates of cell proliferation, in gastric surface epithelial cells in our patients with pernicious anaemia and simple atrophic gastritis. Areas of intestinal metaplasia in gastric mucosa had especially high mitosis counts indicating that such areas may have had a high cell turnover.

The finding of a high gastric DNA-rate in the patient with pernicious anaemia after treatment with vitamin B₁₂ suggests that the rate of production of gastric surface epithelial cells determines, in the resting state, the rate at which these cells are lost from the mucosa.

**SUMMARY**

Using a technique of saline perfusion and continuous suction the rate of accumulation of DNA in the stomach was measured in 28 patients. In 11 the gastric mucosa was normal, nine had pernicious anaemia, and eight simple atrophic gastritis.

The gastric DNA-rate was reproducible (±10%) and was considered to measure cell loss from the gastric mucosa. The mean rate of loss from normal gastric mucosa was calculated to be approximately half a million cells per minute.

Patients with normal gastric mucosa, and with treated pernicious anaemia had similar gastric DNA-rates but a majority of patients with simple atrophic gastritis had much higher values.

In one patient with pernicious anaemia the gastric DNA-rate after treatment with vitamin B₁₂ was three times the rate observed before treatment.

Significantly higher epithelial mitosis counts were found in gastric biopsies of patients with treated pernicious anaemia and simple atrophic gastritis than in those with normal mucosa.

Gastric DNA-rates, mitosis counts, and histological and cytological findings suggested that the gastric DNA-rate was a measure, in the resting state, of gastric epithelial cell production and loss (or turnover); there was a higher than normal turnover of gastric surface epithelial cells in simple atrophic gastritis and in treated pernicious anaemia.

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